Folate and Vitamin B₆ From Diet and Supplements in Relation to Risk of Coronary Heart Disease Among Women

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Context.—Hyperhomocysteinemia is caused by genetic and lifestyle influences, including low intakes of folate and vitamin B₆. However, prospective data relating intake of these vitamins to risk of coronary heart disease (CHD) are not available.

Objective.—To examine intakes of folate and vitamin B₆ in relation to the incidence of nonfatal myocardial infarction (MI) and fatal CHD.

Design.—Prospective cohort study.

Setting and Patients.—In 1980, a total of 80,082 women from the Nurses’ Health Study with no previous history of cardiovascular disease, cancer, hypercholesterolemia, or diabetes completed a detailed food frequency questionnaire from which we derived usual intake of folate and vitamin B₆.

Main Outcome Measure.—Nonfatal MI and fatal CHD confirmed by World Health Organization criteria.

Results.—During 14 years of follow-up, we documented 658 incident cases of nonfatal MI and 281 cases of fatal CHD. After controlling for cardiovascular risk factors, including smoking and hypertension and intake of alcohol, fiber, vitamin E, and saturated, polyunsaturated, and trans fat, the relative risks (RRs) of CHD between extreme quintiles were 0.69 (95% confidence interval [CI], 0.55-0.87) for folate (median intake, 696 µg/d vs 158 µg/d) and 0.67 (95% CI, 0.53-0.85) for vitamin B₆ (median intake, 4.6 mg/d vs 1.1 mg/d). Controlling for the same variables, the RR was 0.55 (95% CI, 0.41-0.74) among women in the highest quintile of both folate and vitamin B₆ intake compared with the opposite extreme. Risk of CHD was reduced among women who regularly used multiple vitamins (RR=0.76; 95% CI, 0.65-0.90), the major source of folate and vitamin B₆, and after excluding multiple vitamin users, among those with higher dietary intakes of folate and vitamin B₆. In a subgroup analysis, compared with nondrinkers, the inverse association between a high-folate diet and CHD was strongest among women who consumed up to 1 alcoholic beverage per day (RR = 0.69; 95% CI, 0.49-0.97) or more than 1 drink per day (RR = 0.27; 95% CI, 0.13-0.58).

Conclusion.—These results suggest that intake of folate and vitamin B₆ above the current recommended dietary allowance may be important in the primary prevention of CHD among women.

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THREE DECADES AGO, premature vascular occlusive disease was identified in patients with inborn metabolic disorders associated with homocysteinuria, leading to the hypothesis that elevated blood homocysteine levels may cause coronary disease. More recently, evidence linking moderately elevated blood homocysteine levels to increased risk has focused attention on genetic and lifestyle determinants of homocysteine levels. Folate and vitamin B₆ are important cofactors for metabolism. Supplementation of the diet above the recommended dietary allowance (RDA) with folate alone, or in combination with vitamin B₁₂, reduces homocysteine levels. The current RDA for folate acid for nonpregnant women is 180 µg/d, and the average dietary intake in this country among adult women is approximately 225 µg/d. Because of evidence that this level of intake may be insufficient to minimize risk of neural tube defects, and possibly coronary heart disease (CHD), some have urged that the RDA be reset to the earlier level of 400 µg/d. 

For editorial comment see p 392.

Although homocysteine may be atherogenic, it also may be only a marker of folate and vitamin B₆ status. Recent epidemiologic evidence suggests that populations with higher plasma levels of folate and pyridoxal phosphate (PLP, the active form of vitamin B₆) have lower risk of carotid artery stenosis and CHD. In prospective study of 130 myocardial infarction (MI) cases and 118 controls, folate intake was inversely associated with CHD risk. To our knowledge, this relation has not been prospectively studied. Furthermore, previous studies have not examined the independent effects of folate and vitamin B₆ from food or from supplements on risk of CHD, nor have previous studies collected sufficient detail to examine subpopulations of individuals at higher risk of CHD due to factors that may directly or indirectly affect circulating levels of folate (eg, smoking, parental history of MI, and alcohol). Therefore, we examined the relation of intakes of folate and vitamin B₆ to risk of CHD among 80,082 women enrolled in the Nurses’ Health Study and followed prospectively for 14 years (1980-1994).
METHODS

The Nurses’ Health Study Cohort

Details of the Nurses’ Health Study have been published elsewhere. Briefly, the cohort was established in 1976 when 121,700 female registered nurses aged 30 to 55 years and residing in 11 large US states completed a mailed questionnaire on their medical history and lifestyle. Every 2 years, follow-up questionnaires have been sent to update information on potential risk factors and identify newly diagnosed cases of CHD and other diseases. In 1980, a 61-item food frequency questionnaire (FFQ) was included to assess intake of micronutrients and other components of diet. In 1984, the FFQ was expanded to include 116 items. Similar questionnaires were used to update diet in 1986 and 1990.

Semiquantitative FFQs

A detailed description of the dietary questionnaires and documentation of their reproducibility and validity have been published elsewhere. Briefly, the questionnaire at baseline contained 61 foods that allowed maximum discrimination of macronutrient and micronutrient intake. The 1980 questionnaire also gathered information on brand of cold cereal, use of vitamin supplements, brand and type of multiple vitamin, usual number taken per week, and years of past supplement use. Refinements to the original instrument led to approximately twice the number of foods on revised questionnaires in 1984, 1986, and 1990. On all dietary questionnaires, a commonly used unit or portion size for each food (eg, 1 egg or 1 slice of bread) was specified, and the participant was asked how often on average during the previous year she had consumed that amount of each food. Nine responses were possible, ranging from “never or less than once per month” to “six or more times per day.” The intake of nutrients was computed by multiplying the frequency of consumption of each unit of food by the nutrient content of the specified portion. Food composition values for folate, riboflavin, vitamin B₆, vitamin B₁₂, methionine, and other nutrients were obtained from the Harvard University Food Composition Database (November 22, 1993) derived from US Department of Agriculture sources and supplemented with manufacturer information. In a detailed validation study, we compared nutrients derived from the 61-item baseline questionnaire with those from four 1-week diet records collected at approximately 3-month intervals and found that the FFQ can provide a reasonable measure of dietary intake of micronutrients among female nurses.

For example, for vitamin B₆, we found a correlation of 0.58 (0.54 for vitamin B₁₂ from foods only) between energy-adjusted nutrient intake derived from the questionnaire and that from the average of four 1-week diet records.

As further validation of the dietary instrument, we, and others, have found that the FFQ predicts circulating levels of folate and vitamin B₁₂. In the Framingham Heart Study, intake calculated from the FFQ predicted plasma folate (r=0.56), plasma vitamin B₁₂ (measured as PLP) (r=0.51), and homocysteine levels.

Population

After up to 4 mailings, 98,462 women returned the 1980 diet questionnaire. A priori, we excluded women who had implausibly high (>14,700 kJ/d [3500 kcal/d]) or low (<2100 kJ/d [500 kcal/d]) total energy intake or those who left 10 or more items blank. We further excluded women with previously diagnosed cancer (n=3526) and cardiovascular disease (ie, angina, MI, stroke, tachycardia, transient ischemic attack; n=1821). In addition, women reporting hypercholesterolemia (n=4122) or diabetes (n=1812) were excluded because these conditions are associated with risk of CHD and these women might tend to change their diet and lifestyle habits after diagnosis. The final 1980 baseline population consisted of 80,082 women.

Outcome

We defined the primary outcome of CHD as documented symptomatic nonfatal MI or fatal CHD occurring after the return of the 1980 questionnaire but before June 1, 1994. We requested permission to review medical records from women who reported a nonfatal MI on a follow-up questionnaire. The records were reviewed by study physicians blinded to the exposure status. We used the World Health Organization criteria for MI; confirmation required symptoms plus either diagnostic electrocardiographic changes or elevated cardiac enzymes. Myocardial infarctions that required hospital admission and for which confirmatory information was obtained by interview or letter, but for which no medical records were available, were designated as “probable.” We included all “confirmed” and “probable” cases in our analyses because results were not substantially different after excluding 17% of cases defined as “probable.”

We were notified of most deaths by the next of kin or the post office. Every 2 years we also search the National Death Index for nonrespondents to biennial questionnaires. We estimate that follow-up for deaths is over 98% complete. Fatal CHD was defined as fatal MI if confirmed by hospital records or autopsy, or if CHD was listed as the primary cause on the death certificate and evidence of previous CHD was available. Information from death certificate alone was not sufficient for the confirmation of CHD.

Data Analysis

All nutrients were energy-adjusted using the residual method. Energy adjustment is based on the a priori biologic considerations that a larger, more physically active person will require a higher caloric intake, which will also be associated with a higher absolute intake of all nutrients. Therefore, by adjusting for energy intake, we examined the composition of the diet, accounting for differences in energy requirements among individuals. The residuals were standardized to the predicted nutrient intake of a woman consuming 6720 kJ/d (1600 kcal/d), the average total energy intake among women responding to the 1980 dietary questionnaire. After energy adjustment, quintiles were created for each nutrient. Person-time for each participant was calculated from the date of return of the 1980 questionnaire to the date of first CHD event, death, or June 1, 1994. Women who reported cardiovascular disease or cancer on previous questionnaires were excluded from subsequent follow-up; thus, each participant could contribute only 1 end point, and the cohort at risk included only those who remained free from a cardiovascular and cancer end point at the beginning of each 2-year follow-up interval.

We calculated incidence rates as the number of events divided by the person-time of follow-up in each quintile. The relative risk (RR) was computed as the rate in a specific quintile of a micronutrient divided by that in the lowest quintile, with adjustment for 5-year age categories. In all multivariate analyses, we calculated the odds ratio of CHD (as an estimate of the RR) for quintiles of baseline (1980) nutrients controlling for age; time period; smoking; body mass index (a measure of weight in kilograms divided by the square of height in meters); postmenopausal hormones; aspirin; vitamin E supplements; exercise; hypertension; parental history of CHD; and intake of polyunsaturated, saturated, and trans fat, fiber, and alcohol. The main analyses were conducted as a pooled logistic regression using baseline nutrients with covariates updated using data from each biennial follow-up. In secondary analyses, incidence of CHD was related to the cumulative average of nutrient intake from dietary questionnaires administered in 1980, 1984, 1986,
and 1990. Incident cases documented between each 2- or 4-year diet questionnaire cycle were examined in relation to the average diet calculated from all the preceding diet measures. Since change in diet after development of intermediate conditions such as angina, hypercholesterolemia, and diabetes may confound the exposure-disease association, we stopped updating diet and covariates at the beginning of the time interval during which individuals developed those intermediate end points.

**RESULTS**

During 14 years of follow-up, we identified 658 incident cases of nonfatal MI and 281 cases of fatal CHD. Based on the assessment of average diet, the largest contributors (in absolute percentage) to the overall intake of folate were multiple vitamins (26%), cold cereal (8%), orange juice (8%), lettuce (7.5%), eggs (3%), broccoli (2.5%), and spinach (2%). Although women with higher folate intake were older and had a higher prevalence of hypertension, they generally weighed less, were less likely to smoke, and ate a diet higher in fiber and lower in saturated fat, polyunsaturated fat, and trans fat (Table 1).

Using the baseline 1980 measure of dietary folate to classify women into quintiles of intake, the age-adjusted RR of CHD was 0.83 (95% confidence interval [CI], 0.43-0.65) comparing women in the highest and lowest quintiles (Table 2). In multivariate models of folate and CHD, smoking and use of vitamin E supplements were the strongest confounders.

The RR for folate was 0.61 (95% CI, 0.49-0.75) after adding smoking to the model and further attenuated (RR=0.69; 95% CI, 0.69-0.79) after controlling for current use of vitamin E supplements. The RR for a diet high in vitamin E supplements (RR=0.75; 95% CI, 0.61-0.92) was slightly stronger than for foods. Although we had substantially fewer cases (n=488) after excluding all women at or below the RDA of 1.6 mg/d, the RR for each 2-mg increase in vitamin E intake was similar when we used the updated cumulative average as a measure of intake rather than only the baseline values and when we used either nonfatal MI or fatal CHD as the outcome.

In more detailed analyses, we examined the independent associations for folate from diet and from supplemental intake. In the same multivariate model noted in Table 2 we simultaneously included folate from both sources; a diet high in folate from food or supplements conferred a reduction in risk of CHD (for food, RR=0.75 [95% CI, 0.68-0.83]; for supplements, RR=0.73 [95% CI, 0.65-0.82]) and quintile 5 (median 2.3 mg/d) vs quintile 1 (median 0.4 mg/d). However, in this model the RR was 0.67 (95% CI, 0.53-0.85) after controlling for measured predictors of CHD (Table 2). In an analysis of the overall trend we found an RR of 0.83 (95% CI, 0.74-0.93) for each 2-mg increase in vitamin E intake. Although we had substantially fewer cases (n=488) after excluding all women at or below the RDA of 1.6 mg/d, the RR for each 2-mg increase in vitamin E intake was similar when we used the updated cumulative average as a measure of intake rather than only the baseline values and when we used either nonfatal MI or fatal CHD as the outcome.

We conducted similar analyses for vitamin B₆ as to those presented above for folate. As with folate, vitamin supplements were by far the largest contributors to total vitamin B₆ intake. Among the top 66% of women not taking a multiple vitamin or other supplements containing vitamin B₆, the leading contributors to intake were beef (14%), cold breakfast cereal (9%), potatoes (9%), bananas (8.5%), chicken (7%), milk (5%), and tuna fish (3.5%). Compared with women in the lowest quintile of vitamin B₆ intake, the age-adjusted RR of CHD was 0.49 (95% CI, 0.40-0.61) among women in the highest quintile. As with folate, the strongest confounders were smoking and use of vitamin E supplements. The final multivariate RR was 0.67 (95% CI, 0.53-0.85) after controlling for measured predictors of CHD (Table 2). In an analysis of the overall trend we found an RR of 0.83 (95% CI, 0.74-0.93) for each 2-mg increase in vitamin B₆ intake. Although we had substantially fewer cases (n=488) after excluding all women at or below the RDA of 1.6 mg/d, the RR for each 2-mg increase in vitamin B₆ intake above the RDA was still suggestive of a reduction in risk of CHD (RR=0.91; 95% CI, 0.79-1.04). In the model that simultaneously assessed the independent associations for vitamin B₆ from food or supplement sources, the inverse association for supplemental intake (RR=0.75; 95% CI, 0.62-0.92; for 2-10 mg/d of supplements vs no supplements) was slightly stronger than for intake from foods (RR=0.85; 95% CI, 0.68-1.07; between quintile 5 [median 2.3 mg/d] and quintile 1 [median 1.1 mg/d]). However, in this model the range in vitamin B₆ intake was much broader for intake of supplements than for foods. The model that incorporated vitamin B₆ assessed from follow-up questionnaires yielded similar results as the multivariate with only baseline dietary measures.

**Table 1.—Baseline (1980) Diet and Lifestyle Characteristics by Quintiles of Total Energy-Adjusted Folate Intake Among 80,082 Women Enrolled in the Nurses’ Health Study**

<table>
<thead>
<tr>
<th>Folate Quintile</th>
<th>1 (&lt;190 µg)</th>
<th>2 (190-244 µg)</th>
<th>3 (245-318 µg)</th>
<th>4 (319-544 µg)</th>
<th>5 (≥545 µg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Folate, µg/d</td>
<td>153</td>
<td>217</td>
<td>278</td>
<td>409</td>
<td>774</td>
</tr>
<tr>
<td>Age, y</td>
<td>44.7</td>
<td>45.7</td>
<td>46.4</td>
<td>46.3</td>
<td>46.3</td>
</tr>
<tr>
<td>Body mass index, kg/m²</td>
<td>24.3</td>
<td>24.4</td>
<td>24.4</td>
<td>24.1</td>
<td>23.9</td>
</tr>
<tr>
<td>Current smokers, %</td>
<td>36.9</td>
<td>29.6</td>
<td>26.4</td>
<td>24.8</td>
<td>24.9</td>
</tr>
<tr>
<td>Multiple vitamins, %</td>
<td>7.7</td>
<td>10.6</td>
<td>14.7</td>
<td>14.3</td>
<td>94.5</td>
</tr>
<tr>
<td>Vitamin E supplements, %</td>
<td>5.9</td>
<td>6.9</td>
<td>8.9</td>
<td>14.3</td>
<td>27.5</td>
</tr>
<tr>
<td>Postmenopausal hormones, %</td>
<td>6.0</td>
<td>5.8</td>
<td>6.3</td>
<td>6.6</td>
<td>7.4</td>
</tr>
<tr>
<td>Regular exercise, %†</td>
<td>35.0</td>
<td>42.2</td>
<td>47.1</td>
<td>50.9</td>
<td>50.6</td>
</tr>
<tr>
<td>Hypertension, %</td>
<td>13.1</td>
<td>14.0</td>
<td>14.7</td>
<td>14.2</td>
<td>14.1</td>
</tr>
<tr>
<td>Parental history of CHD, %‡</td>
<td>19.8</td>
<td>20.0</td>
<td>20.3</td>
<td>19.8</td>
<td>19.7</td>
</tr>
</tbody>
</table>

*Means for each quintile are directly age-standardized using the age distribution of the 80,082 women eligible for analyses.

†Regular exercise is defined as vigorous exercise 1 or more times per week.

‡Parental history of coronary heart disease (CHD) is assigned to women who reported a myocardial infarction in either parent before the age of 65 years.
Table 2.—Relative Risk (RR) and 95% Confidence Intervals (CIs) of Coronary Heart Disease (CHD) (Nonfatal Myocardial Infarction and Fatal CHD) by Quintiles of Total Energy-Adjusted Folate and Vitamin B₆ Intake Among 80,082 Women Enrolled in the Nurses’ Health Study (1980-1994)

<table>
<thead>
<tr>
<th>Quintiles</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>1 1 P for Trend</th>
</tr>
</thead>
<tbody>
<tr>
<td>Folate, median, µg/d</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cases, No.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age-adjusted RR (95% CI)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Multivariate RR (95% CI)*</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vitamin B₆, mg/d (median)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 3.—Relative Risk (RR) and 95% Confidence Interval (CIs) of Coronary Heart Disease (CHD) (Nonfatal Myocardial Infarction and Fatal CHD) for Users of Multiple Vitamin Supplements in 1980 Among 80,082 Women Enrolled in the Nurses’ Health Study (1980-1994)

<table>
<thead>
<tr>
<th>Nonusers</th>
<th>1-3 Pills/wk</th>
<th>4-7 Pills/wk</th>
<th>Current Users*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cases, No.</td>
<td>702</td>
<td>31</td>
<td>184</td>
</tr>
<tr>
<td>Person-years</td>
<td>716,913</td>
<td>42,685</td>
<td>272,435</td>
</tr>
<tr>
<td>Age-adjusted RR</td>
<td>1.0 (Referent)</td>
<td>0.82 (0.57-1.17)</td>
<td>0.87 (0.57-0.79)</td>
</tr>
<tr>
<td>Multivariate RR</td>
<td>1.0 (Referent)</td>
<td>0.97 (0.68-1.40)</td>
<td>0.76 (0.65-0.90)</td>
</tr>
</tbody>
</table>

The reduction in risk associated with a high intake of vitamin B₆ was similar for nonfatal MI or fatal CHD.

COMMENT

During 14 years of follow-up of this large prospective cohort study of women, we found graded associations between higher intakes of folate and vitamin B₆ and lower risk of CHD. For folate, lower risks were seen for higher intake from either food or supplement sources and for intake well above the current RDA of 180 µg/d. For vitamin B₆ the risk was lowest with higher intake of food and supplement sources combined. Risk of CHD was lowest among women with the highest intake of both folate and vitamin B₆. In exploratory analyses of high- and low-risk subgroups, we found the strongest apparent benefit of a high-folate diet among women who consumed alcohol. Our main findings are consistent with the experimental evidence linking higher intake of folate and vitamin B₆ with lower homocysteine levels and with the observational evidence suggesting a strong association between elevated homocysteine levels and cardiovascular disease; however, we cannot exclude the possibility that homocysteine may only be a marker of folate status and other unknown mechanisms could explain the lower rates of CHD among women with higher intakes of folate and vitamin B₆. For example, folate-depleted diets may be prothrombotic.
association with folate could be explained by other healthy lifestyle practices among women in the highest quintile of folate intake. For several variables, such as smoking and use of vitamin E supplements, women in the highest quintile of folate had a lower risk profile (Table 1). Controlling for these factors attenuated the age-adjusted RR from 0.53 (95% CI, 0.43-0.65) to 0.69 (95% CI, 0.53-0.85) (Table 2). However, several lines of evidence suggest that this inverse association for folate is not due to further bias or confounding. First, if the self-selection of the healthiest individuals into the highest quintile of folate were to explain our results, we would expect several other micronutrients found in supplements and “healthy diets” also to be related to lower risk. However, we did not find inverse associations for other such micronutrients (eg, riboflavin, vitamin B12, and vitamin C[13]). Second, our data suggest a graded inverse association between CHD and increasing intake of folate regardless of folate source. Finally, the magnitude of the inverse association for folate is consistent with our parallel study among male health professionals[12] and with the physiologic effects of folate on lowering of homocysteine levels.[3] From calculations by Boushey et al[13] and Tucker et al,[14] increasing folate intake by 100 µg/d would lower homocysteine levels by 6% and risk of CHD by approximately 5%. Each 100 µg/d increase in folate in our population was associated with a 5.8% (95% CI, 2%-9%) lower risk of CHD.

Homocysteine may increase risk of CHD through direct toxicity to endothelial cells, increased coagulation, decreased endothelial reactivity, and stimulation of smooth muscle cell proliferation.[33-36] Higher levels of homocysteine have been observed among patients with peripheral[37,38] and cerebral vascular occlusion[39,40] and coronary disease.[3] Although the positive association is graded between levels of homocysteine and degree of carotid artery stenosis[34] and CHD,[33,35] the association between dietary folate and CHD may not be linear since the association between folate and homocysteine appears to plateau between 400 µg/d and 1 mg/d.[32,33] Within the range of folate intake in this population (10th percentile of 150 µg/d and 90th percentile of 700 µg/d), our results suggest a linear decline in risk of CHD. We were not able to examine benefits of supplementation above 1 mg/d.

Elevated homocysteine levels after a methionine load and fasting homocysteine levels may be independent predictors of CHD.[31,42] Recent evidence suggests that vitamin B6 is generally more effective at lowering fasting homocysteine levels. Our results of a maximum benefit among women in the highest quintile of both folate and vitamin B6 (RR=0.57; 95% CI, 0.40-0.82) are consistent with independent effects of these vitamins in lowering homocysteine levels and suggest that maximum benefit is obtained at optimal levels of both. Randomized trials in secondary and primary prevention are needed to test their independent effects.[43] Although dietary B12 also may lower homocysteine level, plasma B12 levels are influenced more by absorption than intake, thus the lack of association between extreme quintiles of dietary vitamin B12 (RR=1.09; 95% CI, 0.82-1.44) and risk of CHD is expected.

The stronger inverse association between folate and CHD among women who consumed alcohol (Figure) merits further exploration since several subgroup analyses were conducted. At moderate levels of alcohol consumption, oxidation of acetaldehyde from ethanol metabolism can inactivate folate.[44] Furthermore, acetaldehyde binds and inactivates methionine synthase,[45] the enzyme responsible for the folate-dependent remethylation of homocysteine to methionine. Therefore, we would expect that the cardiovascular benefits of moderate alcohol consumption[46] may be partially offset unless women have high folate intake.

In conclusion, in this population of women, higher intakes of folate from food or supplements, alone or in combination with vitamin B6, are associated with substantially lower risk of CHD among women. Our results corroborate and extend recent studies,[14,15] showing lower risk of coronary disease in individuals with higher plasma folate and plasma vitamin B6 levels. Even though in this population of US women average intakes of folate (366 µg/d; median=277 µg/d) and vitamin B6 (3.0 mg/d; median=1.7 mg/d) were well above those from national averages (224 µg/d for folate and 1.51 mg/d for vitamin B6),[11] we still found a graded reduction in risk with higher intake. The lowest risk was among women with intake of folate above 400 µg/d and vitamin B6 above 3 mg/d. Therefore, the current RDA for folate of 180 µg/d and vitamin B6 of 1.6 mg/d deemed sufficient to prevent deficiency among nonpregnant women[15] may not be sufficient to minimize risk of coronary disease. Furthermore, for folate, it is estimated that 88% to 90% of the population has dietary intakes below 400 µg/d[11,23] and that the recently implemented policy of folate supplementation of 140 µg per 100 g of grain products in US food supplies may only increase average intakes by 100 µg/d,[47] and less if currently fortified foods are required to reduce fortification levels.[23] Even after fortification, only 25% of adult women will have dietary folate intake above 400 µg/d. Our results suggest that any widespread increase in folate intake will have a favorable impact on CHD rates, but that maximum benefit would be achieved at folate intake of at least 400 µg/d.

Relative risk of coronary heart disease (nonfatal myocardial infarction and fatal coronary heart disease) by quintiles of energy-adjusted folate across levels of alcohol consumption among 80 082 women in the Nurses’ Health Study. Women in the lowest quintile of folate who did not drink alcohol were the reference category.
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References


